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                 CAplus currency for Korean patents enhanced
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                 comprehensive access to substance and sequence
                  information
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                 to be discontinued
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         SEP 25
                 CA/CAplus current-awareness alert options enhanced
                 to accommodate supplemental CAS indexing of
                 exemplified prophetic substances
NEWS 13
         SEP 26
                 WPIDS, WPINDEX, and WPIX coverage of Chinese and
                 and Korean patents enhanced
NEWS 14
         SEP 29
                 IFICLS enhanced with new super search field
NEWS 15
         SEP 29
                 EMBASE and EMBAL enhanced with new search and
                 display fields
NEWS 16
         SEP 30
                 CAS patent coverage enhanced to include exemplified
                 prophetic substances identified in new Japanese-
                 language patents
                 {\tt EPFULL} enhanced with full implementation of {\tt EPC2000}
NEWS 17
         OCT 07
NEWS 18
         OCT 07 Multiple databases enhanced for more flexible patent
                 number searching
         OCT 22
NEWS 19
                 Current-awareness alert (SDI) setup and editing
                 enhanced
         OCT 22
                 WPIDS, WPINDEX, and WPIX enhanced with Canadian PCT
NEWS 20
                 Applications
NEWS 21
         OCT 24
                 CHEMLIST enhanced with intermediate list of
                 pre-registered REACH substances
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             AND CURRENT DISCOVER FILE IS DATED 23 JUNE 2008.
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chain nodes :

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ring nodes :

1 2 3 4 5 6 7 8 9 10 11 12 13 14 20 21 22 23 24 chain bonds :

ring bonds :

exact/norm bonds :

 $5-7 \quad 6-10 \quad 7-8 \quad 8-9 \quad 8-11 \quad 9-10 \quad 9-14 \quad 11-12 \quad 12-13 \quad 13-14 \quad 15-16 \quad 15-29 \quad 16-17$

18-19 18-31 19-20 20-21 20-24 21-22 22-23 23-24 24-30

exact bonds :

13-15 17-18 17-25 25-26 26-27 26-28

normalized bonds :

1-2 1-6 2-3 3-4 4-5 5-6

Match level:

1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:Atom 8:Atom 9:Atom 10:Atom 11:Atom 12:Atom 13:Atom 14:Atom 15:CLASS 16:CLASS 17:CLASS 18:CLASS 19:CLASS 20:Atom 21:Atom 22:Atom 23:Atom 24:Atom 25:CLASS 26:CLASS 27:CLASS 28:CLASS 29:CLASS 30:CLASS 31:CLASS

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L4 8 L3

 \Rightarrow d 14 1-8 ibib abs hitstr

L4 ANSWER 1 OF 8 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2007:1022710 CAPLUS

DOCUMENT NUMBER: 147:357234

TITLE: Composition containing amidine derivatives or

carboxamide derivatives and steroids, as a medicament

INVENTOR(S): Pignol, Bernadette; Auvin, Serge; Bigg, Dennis;

Chabrier de Lassauniere, Pierre-Etienne

PATENT ASSIGNEE(S): Societe de Conseils de Recherches et d'Applications

Scientifiques (S.C.R.A.S.), Fr.

SOURCE: PCT Int. Appl., 42pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: French

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

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APPLICATION NO.
    PATENT NO.
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    WO 2007101937
                               20070913
                                          WO 2007-FR390
                                                                  20070306
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        W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH,
            CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD,
            GE, GH, GM, GT, HN, HR, HU, ID, IL, IN, IS, JP, KE, KG, KM, KN,
            KP, KR, KZ, LA, LC, LK, LR, LS, LT, LU, LY, MA, MD, MG, MK, MN,
            MW, MX, MY, MZ, NA, NG, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RS,
            RU, SC, SD, SE, SG, SK, SL, SM, SV, SY, TJ, TM, TN, TR, TT, TZ,
            UA, UG, US, UZ, VC, VN, ZA, ZM, ZW
         RW: AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE,
            IS, IT, LT, LU, LV, MC, MT, NL, PL, PT, RO, SE, SI, SK, TR, BF,
            BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG, BW,
            GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ,
            BY, KG, KZ, MD, RU, TJ, TM
                               20070914
                                           FR 2006-2000
                                                                  20060307
    FR 2898274
                         Α1
                               20081003
    FR 2898274
                         В1
PRIORITY APPLN. INFO.:
                                           FR 2006-2000
                                                               A 20060307
                        MARPAT 147:357234
OTHER SOURCE(S):
    The present invention relates to a composition containing at least one amidine
    derivative or carboxamide derivative (Markush included) in combination with at
    least one compound chosen from steroids, corticoids or corticosteroids,
    wherein the composition is suitable for the preparation of a medicament.
Compound
    preparation is included.
     339007-48-6 339007-48-6D, salts 339007-76-0
ΙT
    339007-76-0D, salts 742104-24-1 742104-24-1D,
    salts 866006-13-5 866006-13-5D, salts
    866006-14-6 866006-14-6D, salts 866006-16-8
    866006-16-8D, salts 866006-17-9 866006-17-9D,
    salts
    RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL
     (Biological study); USES (Uses)
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(amidine derivative or carboxamide derivative combination with steroid for

(acetyloxy)tetrahydro-3-furanyl]amino]carbonyl]-3-methylbutyl]- (CA INDEX

KIND

DATE

DATE

Absolute stereochemistry.

NAME)

RN

CN

therapeutic)

339007-48-6 CAPLUS

339007-48-6 CAPLUS RN CN 10H-Phenothiazine-2-carboxamide, N-[(1S)-1-[[(2S,3S)-2-(acetyloxy)tetrahydro-3-furanyl]amino]carbonyl]-3-methylbutyl]- (CA INDEX NAME)

10H-Phenothiazine-2-carboxamide, N-[(1S)-1-[[(2S,3S)-2-

RN 339007-76-0 CAPLUS

CN 10H-Phenothiazine-2-carboxamide, N-[(1S)-1-[[[(2R,3S)-2-(acetyloxy)tetrahydro-3-furanyl]amino]carbonyl]-3-methylbutyl]- (CA INDEX NAME)

Absolute stereochemistry.

RN 339007-76-0 CAPLUS

CN 10H-Phenothiazine-2-carboxamide, N-[(1S)-1-[[(2R,3S)-2-(acetyloxy)tetrahydro-3-furanyl]amino]carbonyl]-3-methylbutyl]- (CA INDEX NAME)

Absolute stereochemistry.

RN 742104-24-1 CAPLUS

CN 10H-Phenothiazine-2-carboxamide, N-[(1S)-1-[[[(3S)-2-(acetyloxy)tetrahydro-3-furanyl]amino]carbonyl]-3-methylbutyl]- (CA INDEX NAME)

Absolute stereochemistry.

RN 742104-24-1 CAPLUS

CN 10H-Phenothiazine-2-carboxamide, N-[(1S)-1-[[[(3S)-2-(acetyloxy)tetrahydro-3-furanyl]amino]carbonyl]-3-methylbutyl]- (CA INDEX NAME)

RN 866006-13-5 CAPLUS

CN 10H-Phenothiazine-2-carboxamide, N-[(1S)-3-methyl-1-[[[(2S,3S)-tetrahydro-2-hydroxy-3-furanyl]amino]carbonyl]butyl]- (CA INDEX NAME)

Absolute stereochemistry.

RN 866006-13-5 CAPLUS

CN 10H-Phenothiazine-2-carboxamide, N-[(1S)-3-methyl-1-[[[(2S,3S)-tetrahydro-2-hydroxy-3-furanyl]amino]carbonyl]butyl]- (CA INDEX NAME)

Absolute stereochemistry.

RN 866006-14-6 CAPLUS

CN 10H-Phenothiazine-2-carboxamide, N-[(1S)-3-methyl-1-[[[(2R,3S)-tetrahydro-2-hydroxy-3-furanyl]amino]carbonyl]butyl]- (CA INDEX NAME)

Absolute stereochemistry.

RN 866006-14-6 CAPLUS

CN 10H-Phenothiazine-2-carboxamide, N-[(1S)-3-methyl-1-[[[(2R,3S)-tetrahydro-2-hydroxy-3-furanyl]amino]carbonyl]butyl]- (CA INDEX NAME)

RN 866006-16-8 CAPLUS

CN 10H-Phenothiazine-2-carboxamide, N-[(1R)-3-methyl-1-[[[(2S,3S)-tetrahydro-2-hydroxy-3-furanyl]amino]carbonyl]butyl]- (CA INDEX NAME)

Absolute stereochemistry.

RN 866006-16-8 CAPLUS

CN 10H-Phenothiazine-2-carboxamide, N-[(1R)-3-methyl-1-[[[(2S,3S)-tetrahydro-2-hydroxy-3-furanyl]amino]carbonyl]butyl]- (CA INDEX NAME)

Absolute stereochemistry.

RN 866006-17-9 CAPLUS

CN 10H-Phenothiazine-2-carboxamide, N-[(1R)-3-methyl-1-[[[(2R,3S)-tetrahydro-2-hydroxy-3-furanyl]amino]carbonyl]butyl]- (CA INDEX NAME)

Absolute stereochemistry.

RN 866006-17-9 CAPLUS

CN 10H-Phenothiazine-2-carboxamide, N-[(1R)-3-methyl-1-[[[(2R,3S)-tetrahydro-2-hydroxy-3-furanyl]amino]carbonyl]butyl]- (CA INDEX NAME)

REFERENCE COUNT: 5 THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 2 OF 8 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2007:452302 CAPLUS

DOCUMENT NUMBER: 147:64455

TITLE: A novel dual inhibitor of calpains and lipid

peroxidation (BN82270) rescues the cochlea from sound

trauma

AUTHOR(S): Wang, Jing; Pignol, Bernadette; Chabrier,

Pierre-Etienne; Saido, Takaomi; Lloyd, Ruth; Tang,

Yong; Lenoir, Marc; Puel, Jean-Luc

CORPORATE SOURCE: Laboratoire de Physiopathologie et Therapie des

Deficits Sensoriels et Moteurs, INSERM U583,

Montpellier, Fr.

SOURCE: Neuropharmacology (2007), 52(6), 1426-1437

CODEN: NEPHBW; ISSN: 0028-3908

PUBLISHER: Elsevier B.V.

DOCUMENT TYPE: Journal LANGUAGE: English

AB Free radical and calcium buffering mechanisms are implicated in cochlear cell damage that has been induced by sound trauma. Thus in this study we evaluated the therapeutic effect of a novel dual inhibitor of calpains and of lipid peroxidn. (BN 82270) on the permanent hearing and hair cell loss induced by sound trauma. Perfusion of BN 82270 into the scala tympani of the guinea pig cochlea prevented the formation of calpain-cleaved fodrin, translocation of cytochrome c, DNA fragmentation and hair cell degeneration caused by sound trauma. This was confirmed by functional tests in vivo, showing a clear dose-dependent reduction of permanent hearing loss (ED50 = 4.07 μM) with almost complete protection at 100 μM . Furthermore, BN82270 still remained effective even when applied onto the round window membrane after sound trauma had occurred, within a therapeutic window of 24 h. This indicates that BN 82270 may be of potential therapeutic value in treating the cochlea after sound trauma.

IT 742104-24-1, BN 82270

RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(novel dual inhibitor of calpains and lipid peroxidn. (BN82270) rescues the cochlea from sound trauma)

RN 742104-24-1 CAPLUS

CN 10H-Phenothiazine-2-carboxamide, N-[(1S)-1-[[[(3S)-2-(acetyloxy)tetrahydro-3-furanyl]amino]carbonyl]-3-methylbutyl]- (CA INDEX NAME)

REFERENCE COUNT: 96 THERE ARE 96 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 3 OF 8 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2007:325817 CAPLUS

DOCUMENT NUMBER: 146:372657

TITLE: Calpain inhibitors and antioxidants act

synergistically to prevent cell necrosis: effects of the novel dual inhibitors (cysteine protease inhibitor and antioxidant) BN 82204 and its pro-drug BN 82270.

[Erratum to document cited in CA146:075148]

AUTHOR(S): Pignol, Bernadette; Auvin, Serge; Carre, Denis; Marin,

Jean-Gregoire; Chabrier, Pierre-Etienne

CORPORATE SOURCE: Department of Neurobiology, Ipsen Research

Laboratories, Les Ulis, Fr.

SOURCE: Journal of Neurochemistry (2007), 100(5), 1430

CODEN: JONRA9; ISSN: 0022-3042

PUBLISHER: Blackwell Publishing Ltd.

DOCUMENT TYPE: Journal LANGUAGE: English

AB On page 1224, right column, second full paragraph, line 2, "BHT-PD150606" should read: "BHT+PD150606". On page 1224, right column, second full paragraph, line 2, "BHT++calpeptin" should read: "BHT+calpeptin". On page 1224, right column, second full paragraph, line 3,

"4-hydroxydiphenylamine++Z-Leu-Leu-H" should read:

"4-hydroxydiphenylamine+Z-Leu-Leu-H". On page 1224, right column, second full paragraph, line 3, "BHT-Z-Leu-Leu-H" should read: "BHT+Z-Leu-Leu-H".

IT 339007-47-5, BN 82204 742104-24-1, BN 82270

RL: DMA (Drug mechanism of action); PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(use of calpain inhibitors and antioxidants in combination or single BN 82204 and its pro-drug BN 82270 with multiple activities to prevent cell necrosis (Erratum))

RN 339007-47-5 CAPLUS

CN 10H-Phenothiazine-2-carboxamide, N-[(1S)-3-methyl-1-[[[(3S)-tetrahydro-2-hydroxy-3-furanyl]amino]carbonyl]butyl]- (CA INDEX NAME)

Absolute stereochemistry.

RN 742104-24-1 CAPLUS

CN 10H-Phenothiazine-2-carboxamide, N-[(1S)-1-[[[(3S)-2-(acetyloxy)tetrahydro-3-furanyl]amino]carbonyl]-3-methylbutyl]- (CA INDEX NAME)

Absolute stereochemistry.

REFERENCE COUNT: THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS 1 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 4 OF 8 CAPLUS COPYRIGHT 2008 ACS on STN L4

ACCESSION NUMBER: 2006:1135981 CAPLUS

DOCUMENT NUMBER: 146:75148

Calpain inhibitors and antioxidants act TITLE:

> synergistically to prevent cell necrosis: effects of the novel dual inhibitors (cysteine protease inhibitor and antioxidant) BN 82204 and its pro-drug BN 82270

AUTHOR(S): Pignol, Bernadette; Auvin, Serge; Carre, Denis; Marin,

Jean-Gregoire; Chabrier, Pierre-Etienne

CORPORATE SOURCE: Department of Neurobiology, Ipsen Research

Laboratories, Les Ulis, Fr.

Journal of Neurochemistry (2006), 98(4), 1217-1228 SOURCE:

CODEN: JONRA9; ISSN: 0022-3042

PUBLISHER: Blackwell Publishing Ltd.

DOCUMENT TYPE: Journal LANGUAGE: English

Cell death is a common feature observed in neurodegenerative disorders, and is often associated with calpain activation and overprodn. of reactive oxygen species (ROS). This study investigated the use of calpain inhibitors and antioxidants in combination to protect cells against necrosis. Maitotoxin (MTX), which induces a massive influx of calcium, was used to provoke neuronal cell death. This toxin increased, in a concentration-dependent manner,

both calpain activity and ROS formation. Calpain inhibitors or antioxidants inhibited MTX-induced necrosis only marginally (below 20%), whereas their association protected against cell death by 40-66% in a synergistic manner. BN 82204, which possesses both calpain-cathepsin L inhibitory and antioxidant properties, and its acetylated pro-drug BN 82270, totally protected cells at 100 μM . The pro-drug BN 82270, which had better cell penetration, was twice as effective as the active principle BN 82204 in protecting glioma C6 or neuroblastoma SHSY5Y cells against death. These results suggest the potential therapeutic relevance of using a single mol. with multiple activities (cysteine protease inhibitor/antioxidant), and warrant further in vivo investigations in models of neuronal disorders.

339007-47-5, BN 82204 742104-24-1, BN 82270 IT

> RL: DMA (Drug mechanism of action); PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(use of calpain inhibitors and antioxidants in combination or single BN 82204 and its pro-drug BN 82270 with multiple activities to prevent cell necrosis)

339007-47-5 CAPLUS RN

CN hydroxy-3-furanyl]amino]carbonyl]butyl]- (CA INDEX NAME)

RN 742104-24-1 CAPLUS

CN 10H-Phenothiazine-2-carboxamide, N-[(1S)-1-[[[(3S)-2-(acetyloxy)tetrahydro-3-furanyl]amino]carbonyl]-3-methylbutyl]- (CA INDEX NAME)

Absolute stereochemistry.

REFERENCE COUNT: 43 THERE ARE 43 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 5 OF 8 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2006:583794 CAPLUS

DOCUMENT NUMBER: 145:116788

TITLE: Treatment of rats with calpain inhibitors prevents

sepsis-induced muscle proteolysis independent of

atrogin-1/MAFbx and MuRF1 expression

AUTHOR(S): Fareed, Moin U.; Evenson, Amy R.; Wei, Wei; Menconi,

Michael; Poylin, Vitaliy; Petkova, Victoria; Pignol,

Bernadette; Hasselgren, Per-Olof

CORPORATE SOURCE: Department of Surgery, Harvard Medical School, Boston,

MA, USA

SOURCE: American Journal of Physiology (2006), 290(6, Pt. 2),

R1589-R1597

CODEN: AJPHAP; ISSN: 0002-9513
PUBLISHER: American Physiological Society

DOCUMENT TYPE: Journal LANGUAGE: English

Muscle wasting in sepsis is a significant clin. problem because it results AΒ in muscle weakness and fatigue that may delay ambulation and increase the risk for thromboembolic and pulmonary complications. Treatments aimed at preventing or reducing muscle wasting in sepsis, therefore, may have important clin. implications. Recent studies suggest that sepsis-induced muscle proteolysis may be initiated by calpain-dependent release of myofilaments from the sarcomere, followed by ubiquitination and degradation of the myofilaments by the 26S proteasome. In the present expts., treatment of rats with one of the calpain inhibitors calpeptin or BN82270 inhibited protein breakdown in muscles from rats made septic by cecal ligation and puncture. The inhibition of protein breakdown was not accompanied by reduced expression of the ubiquitin ligases atrogin-1/MAFbx and MuRF1, suggesting that the ubiquitin-proteasome system is regulated independent of the calpain system in septic muscle. When incubated muscles were treated in vitro with calpain inhibitor, protein breakdown rates and calpain activity were reduced, consistent with a direct effect in skeletal muscle. Addnl. expts. suggested that the effects of BN82270 on muscle

protein breakdown may, in part, reflect inhibited cathepsin L activity, in addition to inhibited calpain activity. When cultured myoblasts were transfected with a plasmid expressing the endogenous calpain inhibitor calpastatin, the increased protein breakdown rates in dexamethasone-treated myoblasts were reduced, supporting a role of calpain activity in atrophying muscle. The present results suggest that treatment with calpain inhibitors may prevent sepsis-induced muscle wasting. 742104-24-1, BN 82270

RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(treatment of rats with calpain inhibitors prevents sepsis-induced muscle proteolysis independent of atrogin-1/MAFbx and MuRF1 expression) 742104-24-1 CAPLUS

RN 742104-24-1 CAPLUS CN 10H-Phenothiazine-2-

TT

N 10H-Phenothiazine-2-carboxamide, N-[(1S)-1-[[[(3S)-2-(acetyloxy)tetrahydro-3-furanyl]amino]carbonyl]-3-methylbutyl]- (CA INDEX NAME)

Absolute stereochemistry.

REFERENCE COUNT: 46 THERE ARE 46 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 6 OF 8 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2005:1075636 CAPLUS

DOCUMENT NUMBER: 143:339689

TITLE: Use of a phenothiazine derivative for preventing

and/or treating hearing loss

INVENTOR(S): Pignol, Bernadette; Puel, Jean-Luc; Auvin, Serge;

Chabrier de Lassauniere, Pierre-Etienne; Wang, Jing

PATENT ASSIGNEE(S): Societe de Conseils de Recherches et d'Applications

Scientifiques S.C.R.A.S., Fr.

SOURCE: PCT Int. Appl., 33 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: French

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

| PA | CENT | NO. | | | KIND | | DATE | | | APPLICATION NO. | | | | | DATE | | | |
|------------|------------|------|-----|-----|------|----------|------|------|-----|-----------------|------|----------|-----|----------|----------|-----|-----|----|
| WO | 2005 | 0923 | 45 | | A1 | | 2005 | 1006 | | WO 2 | 005- | FR71 | | 20050325 | | | | |
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| | | CN, | CO, | CR, | CU, | CZ, | DE, | DK, | DM, | DZ, | EC, | EE, | EG, | ES, | FΙ, | GB, | GD, | |
| | | GE, | GH, | GM, | HR, | HU, | ID, | IL, | IN, | IS, | JP, | ΚE, | KG, | KP, | KR, | KΖ, | LC, | |
| | | LK, | LR, | LS, | LT, | LU, | LV, | MA, | MD, | MG, | MK, | MN, | MW, | MX, | MZ, | NA, | NI, | |
| | | NO, | NZ, | OM, | PG, | PH, | PL, | PT, | RO, | RU, | SC, | SD, | SE, | SG, | SK, | SL, | SM, | |
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| | RW: | BW, | GH, | GM, | KE, | LS, | MW, | MZ, | NA, | SD, | SL, | SZ, | TZ, | UG, | ZM, | ZW, | AM, | |
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| | | EE, | ES, | FΙ, | FR, | GB, | GR, | HU, | ΙE, | IS, | IT, | LT, | LU, | MC, | NL, | PL, | PT, | |
| | | RO, | SE, | SI, | SK, | TR, | BF, | ВJ, | CF, | CG, | CI, | CM, | GA, | GN, | GQ, | G₩, | ML, | |
| | | MR, | NE, | SN, | TD, | TG | | | | | | | | | | | | |
| FR | FR 2867979 | | | | A1 | 20050930 | | | | FR 2004-3203 | | | | | 20040329 | | | |
| FR 2867979 | | | | | В1 | | 2006 | 0630 | | | | | | | | | | |

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CA 2560988
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     EP 1732567
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                           Α1
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PRIORITY APPLN. INFO.:
                                             FR 2004-3203
                                                                    20040329
                                             FR 2004-6404
                                                                     20040614
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                                             WO 2005-FR713
                                                                  W
                                                                     20050325
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OTHER SOURCE(S): MARPAT 143:339689

GΙ

AB The invention discloses the use of a phenothiazine derivative I (R = H, alkyl, aralkyl, etc.) for preparing a medicine for preventing and/or treating hearing loss.

Ι

IT 339007-47-5 339007-47-5D, derivs. 742104-24-1
866006-13-5 866006-13-5D, derivs. 866006-14-6
866006-14-6D, derivs. 866006-15-7 866006-15-7D
, derivs. 866006-16-8 866006-16-8D, derivs.
866006-17-9 866006-17-9D, derivs.
RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL

(Biological study); USES (Uses)

(phenothiazine derivs. for prevention and/or treatment of hearing loss) 339007-47-5 CAPLUS

RN 339007-47-5 CAPLUS
CN 10H-Phenothiazine-2-carboxamide, N-[(1S)-3-methyl-1-[[[(3S)-tetrahydro-2-hydroxy-3-furanyl]amino]carbonyl]butyl]- (CA INDEX NAME)

Absolute stereochemistry.

RN 339007-47-5 CAPLUS

CN 10H-Phenothiazine-2-carboxamide, N-[(1S)-3-methyl-1-[[[(3S)-tetrahydro-2-hydroxy-3-furanyl]amino]carbonyl]butyl]- (CA INDEX NAME)

RN 742104-24-1 CAPLUS

CN 10H-Phenothiazine-2-carboxamide, N-[(1S)-1-[[[(3S)-2-(acetyloxy)tetrahydro-3-furanyl]amino]carbonyl]-3-methylbutyl]- (CA INDEX NAME)

Absolute stereochemistry.

RN 866006-13-5 CAPLUS

CN 10H-Phenothiazine-2-carboxamide, N-[(1S)-3-methyl-1-[[[(2S,3S)-tetrahydro-2-hydroxy-3-furanyl]amino]carbonyl]butyl]- (CA INDEX NAME)

Absolute stereochemistry.

RN 866006-13-5 CAPLUS

CN 10H-Phenothiazine-2-carboxamide, N-[(1S)-3-methyl-1-[[[(2S,3S)-tetrahydro-2-hydroxy-3-furanyl]amino]carbonyl]butyl]- (CA INDEX NAME)

Absolute stereochemistry.

RN 866006-14-6 CAPLUS

CN 10H-Phenothiazine-2-carboxamide, N-[(1S)-3-methyl-1-[[[(2R,3S)-tetrahydro-2-hydroxy-3-furanyl]amino]carbonyl]butyl]- (CA INDEX NAME)

RN 866006-14-6 CAPLUS

CN 10H-Phenothiazine-2-carboxamide, N-[(1S)-3-methyl-1-[[[(2R,3S)-tetrahydro-2-hydroxy-3-furanyl]amino]carbonyl]butyl]- (CA INDEX NAME)

Absolute stereochemistry.

RN 866006-15-7 CAPLUS

CN 10H-Phenothiazine-2-carboxamide, N-[(1R)-3-methyl-1-[[[(3S)-tetrahydro-2-hydroxy-3-furanyl]amino]carbonyl]butyl]- (CA INDEX NAME)

Absolute stereochemistry.

RN 866006-15-7 CAPLUS

CN 10H-Phenothiazine-2-carboxamide, N-[(1R)-3-methyl-1-[[[(3S)-tetrahydro-2-hydroxy-3-furanyl]amino]carbonyl]butyl]- (CA INDEX NAME)

Absolute stereochemistry.

RN 866006-16-8 CAPLUS

CN 10H-Phenothiazine-2-carboxamide, N-[(1R)-3-methyl-1-[[[(2S,3S)-tetrahydro-2-hydroxy-3-furanyl]amino]carbonyl]butyl]- (CA INDEX NAME)

RN 866006-16-8 CAPLUS

CN 10H-Phenothiazine-2-carboxamide, N-[(1R)-3-methyl-1-[[[(2S,3S)-tetrahydro-2-hydroxy-3-furanyl]amino]carbonyl]butyl]- (CA INDEX NAME)

Absolute stereochemistry.

RN 866006-17-9 CAPLUS

CN 10H-Phenothiazine-2-carboxamide, N-[(1R)-3-methyl-1-[[[(2R,3S)-tetrahydro-2-hydroxy-3-furanyl]amino]carbonyl]butyl]- (CA INDEX NAME)

Absolute stereochemistry.

RN 866006-17-9 CAPLUS

CN 10H-Phenothiazine-2-carboxamide, N-[(1R)-3-methyl-1-[[[(2R,3S)-tetrahydro-2-hydroxy-3-furanyl]amino]carbonyl]butyl]- (CA INDEX NAME)

Absolute stereochemistry.

REFERENCE COUNT: 10 THERE ARE 10 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 7 OF 8 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2004:498592 CAPLUS

DOCUMENT NUMBER: 141:207514

TITLE: Novel dual inhibitors of calpain and lipid

peroxidation

AUTHOR(S): Auvin, Serge; Pignol, Bernadette; Navet, Edith; Pons,

Dominique; Marin, Jean-G.; Bigg, Dennis; Chabrier,

Pierre-E.

CORPORATE SOURCE: Department of Medicinal Chemistry, Ipsen Research

Laboratories, Institut Henri Beaufour, Les Ulis,

91966, Fr.

SOURCE: Bioorganic & Medicinal Chemistry Letters (2004),

14(14), 3825-3828

CODEN: BMCLE8; ISSN: 0960-894X

PUBLISHER: Elsevier Science B.V.

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 141:207514

GΙ

AB A series of mols. I (R1 = phenothiazin-1-yl, phenothiazin-2-yl, 1-benzyl-5-indolinylamino, etc., R2 = H; R1 = phenothiazin-2-yl, R2 = MeCO) with dual inhibitory activities on calpain and lipid peroxidn. were synthesized. These hybrid compds. were built on the calpain pharmacophore 2-hydroxytetrahydrofuran linked to a set of antioxidants via a L-leucine linker. I (R1 = phenothiazin-2-yl, R2 = MeCO), the most potent in cellular calpain and lipid peroxidn. inhibitions, provided effective protection against glial cell death induced by maitotoxin.

IT 339007-47-5P

RL: PAC (Pharmacological activity); RCT (Reactant); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); RACT (Reactant or reagent)

(preparation of (peptidyl)(hydroxy)tetrahydrofurans as dual inhibitors of calpain and lipid peroxidn.)

RN 339007-47-5 CAPLUS

CN 10H-Phenothiazine-2-carboxamide, N-[(1S)-3-methyl-1-[[[(3S)-tetrahydro-2-hydroxy-3-furanyl]amino]carbonyl]butyl]- (CA INDEX NAME)

Absolute stereochemistry.

IT 742104-24-1P, BN 82270

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)

(preparation of (peptidyl)(hydroxy)tetrahydrofurans as dual inhibitors of calpain and lipid peroxidn.)

RN 742104-24-1 CAPLUS

CN 10H-Phenothiazine-2-carboxamide, N-[(1S)-1-[[[(3S)-2-(acetyloxy)tetrahydro-

3-furanyl]amino]carbonyl]-3-methylbutyl]- (CA INDEX NAME)

Absolute stereochemistry.

REFERENCE COUNT: 15 THERE ARE 15 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 8 OF 8 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2001:338525 CAPLUS

DOCUMENT NUMBER: 134:353248

TITLE: Novel heterocyclic compounds and their use as

medicines

INVENTOR(S): Auvin, Serge; Chabrier De Lassauniere, Pierre-Etienne PATENT ASSIGNEE(S):

Societe De Conseils De Recherches Et D'applications

Scientifiques (S.C.R.A.S.), Fr. SOURCE:

PCT Int. Appl., 77 pp. CODEN: PIXXD2

DOCUMENT TYPE: Patent

French LANGUAGE:

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

| PA: | FENT | NO. | | | KIN | | DATE | | APPLICATION NO. | | | | | | DATE | | | |
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| | | | | | A2 20010510 A3 20010927 | | | , | WO 2 | 000- | | 20001103 | | | | | | |
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| | 0000 | | | | | | GA, | , | , | , | , | , | , | , | | 0001 | 405 | |
| | FR 2800737 FR 2800737 | | | | | | 2001 | | | FR 1 | 999- | 1385 | 8 | | 1 | 9991 | 105 | |
| | | | | | 2006 | | | DD 3 | 000 | CE 2 E | | | 2 | 0000 | E 0 0 | | | |
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| | 1233 | | - | | | | 2002 | | | | | | | | | | | |
| | 1233 | | | | | | 2002 | | | UL 2 | 000 | <i>J</i> 1 ± 0 | 10 | | | 0001 | 103 | |
| ш | | | | | | | ES, | | GB. | GR. | TT. | LT. | LII. | NI. | SE. | MC - | PT. | |
| | | | | | | | RO, | | | | | , | шо, | 1111, | 01, | 110, | , | |
| HU | 2002 | | , | , | , | | 2003 | , | | , | | 3183 | | | 2 | 0001 | 103 | |
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| JР | 2003 | 5130 | | | | | 2003 | | | JP 2 | 001- | 5348 | 05 | | 2 | 0001 | 103 | |
| ΝZ | 5184 | 518420 | | | | | 2004 | 0227 | | NZ 2 | 000- | 5184 | 20 | | 2 | 0001 | 103 | |
| ΑU | 7815 | 51 | | | В2 | | 2005 | 0526 | | AU 2 | 001- | 1287 | 1 | | 2 | 0001 | 103 | |
| RU | 2260 | | | | | | 20050910 RU 2002-114696 | | | | | 20001103 | | | | | | |
| AT | 3188 | 09 | | | T | | 2006 | 0315 | | AT 2 | 000- | 9746 | 46 | | 2 | 0001 | 103 | |
| EP | 1661 | | | | A1 | | 2006 | | | EP 2 | 005- | 7719 | 4 | | 2 | 0001 | 103 | |

R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, FI, CY, TR PT 2000-974646 PT 1233962 Т 20060731 20001103 ES 2259617 ES 2000-974646 20001103 T3 20061016 CN 1317278 С 20070523 CN 2000-815926 20001103 US 6747024 В1 20040608 US 2002-111994 20020430 NO 2002002088 Α 20020502 NO 2002-2088 20020502 MX 2002PA04442 Α 20020902 MX 2002-PA4442 20020503 IN 2002MN00604 20050304 IN 2002-MN604 20020513 Α HK 1052706 20070928 HK 2003-105058 A1 20030714 US 20040180936 20040916 US 2004-803387 Α1 20040316 AU 2005-203713 AU 2005203713 Α1 20050915 20050818 PRIORITY APPLN. INFO.: FR 1999-13858 19991105 Α FR 2000-6535 A 20000523 EP 2000-974646 A3 20001103 WO 2000-FR3067 W 20001103 US 2002-111994 A3 20020430

OTHER SOURCE(S): MARPAT 134:353248

AB Novel heterocyclic derivs. which have calpain inhibiting and/or reactive oxygen species trapping activity (no data) are reported. Thus, (R)-Trolox was treated with (S)-2-aminobutyrolactone hydrochloride, followed by DIBAL reduction to give (2R)-6-hydroxy-N-[(3S)-2-hydroxytetrahydrofuran-3-yl]-2,5,7,8-tetramethyl-3,4-dihydro-2H-chromene-2-carboxamide.

IT 339007-47-5P 339007-48-6P 339007-52-2P 339007-53-3P 339007-54-4P 339007-55-5P 339007-56-6P 339007-57-7P 339007-76-0P

RL: SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of novel heterocyclic compds. as calpain inhibitors and trapping agents for reactive oxygen species)

RN 339007-47-5 CAPLUS

CN 10H-Phenothiazine-2-carboxamide, N-[(1S)-3-methyl-1-[[[(3S)-tetrahydro-2-hydroxy-3-furanyl]amino]carbonyl]butyl]- (CA INDEX NAME)

Absolute stereochemistry.

RN 339007-48-6 CAPLUS

CN 10H-Phenothiazine-2-carboxamide, N-[(1S)-1-[[[(2S,3S)-2-(acetyloxy)tetrahydro-3-furanyl]amino]carbonyl]-3-methylbutyl]- (CA INDEX NAME)

Absolute stereochemistry.

RN 339007-52-2 CAPLUS

CN Propanoic acid, 2,2-dimethyl-, (3S)-tetrahydro-3-[[(2S)-4-methyl-1-oxo-2-[(10H-phenothiazin-2-ylcarbonyl)amino]pentyl]amino]-2-furanyl ester (CA INDEX NAME)

Absolute stereochemistry.

RN 339007-53-3 CAPLUS

CN Butanoic acid, 3,3-dimethyl-, (3S)-tetrahydro-3-[[(2S)-4-methyl-1-oxo-2-[(10H-phenothiazin-2-ylcarbonyl)amino]pentyl]amino]-2-furanyl ester (CA INDEX NAME)

Absolute stereochemistry.

RN 339007-54-4 CAPLUS

CN 10H-Phenothiazine-2-carboxamide, N-[(1S)-1-[[[(3S)-2-(benzoyloxy)tetrahydro-3-furanyl]amino]carbonyl]-3-methylbutyl]- (CA INDEX NAME)

Absolute stereochemistry.

RN 339007-55-5 CAPLUS

CN Benzeneacetic acid, (3S)-tetrahydro-3-[[(2S)-4-methyl-1-oxo-2-[(10H-phenothiazin-2-ylcarbonyl)amino]pentyl]amino]-2-furanyl ester (CA INDEX NAME)

RN 339007-56-6 CAPLUS

CN L-Phenylalanine, N,N-dimethyl-, (3S)-tetrahydro-3-[[(2S)-4-methyl-1-oxo-2-[(10H-phenothiazin-2-ylcarbonyl)amino]pentyl]amino]-2-furanyl ester (CA INDEX NAME)

Absolute stereochemistry.

RN 339007-57-7 CAPLUS

CN 4-Morpholinecarboxylic acid, (3S)-tetrahydro-3-[[(2S)-4-methyl-1-oxo-2-[(10H-phenothiazin-2-ylcarbonyl)amino]pentyl]amino]-2-furanyl ester (CA INDEX NAME)

Absolute stereochemistry.

RN 339007-76-0 CAPLUS

CN 10H-Phenothiazine-2-carboxamide, N-[(1S)-1-[[[(2R,3S)-2-(acetyloxy)tetrahydro-3-furanyl]amino]carbonyl]-3-methylbutyl]- (CA INDEX NAME)

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NEWS 12 FEB 02 GENBANK enhanced with SET PLURALS and SET SPELLING

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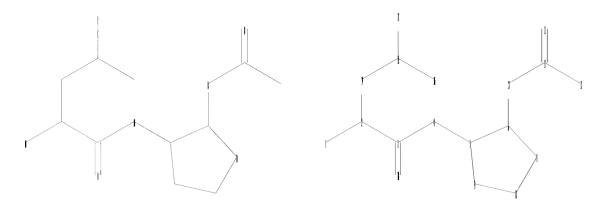
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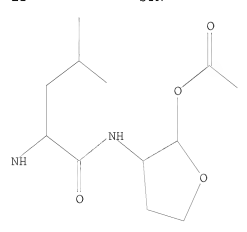
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ring bonds :
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Match level :

1:CLASS 2:CLASS 3:CLASS 4:CLASS 5:Atom 6:Atom 7:Atom 8:Atom 9:Atom 10:CLASS 11:CLASS 12:CLASS 13:CLASS 14:CLASS 15:CLASS 16:CLASS 17:CLASS 18:CLASS

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SAMPLE SCREEN SEARCH COMPLETED - 9 TO ITERATE

100.0% PROCESSED 9 ITERATIONS 5 ANSWERS

SEARCH TIME: 00.00.02

FULL FILE PROJECTIONS: ONLINE **COMPLETE**
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PROJECTED ITERATIONS: 9 TO 360 PROJECTED ANSWERS: 5 TO 234

L2 5 SEA SSS SAM L1

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100.0% PROCESSED 157 ITERATIONS 88 ANSWERS

SEARCH TIME: 00.00.01

L3 88 SEA SSS FUL L1

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SINCE FILE TOTAL
ENTRY SESSION
FULL ESTIMATED COST
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186.10

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FILE COVERS 1907 - 4 Feb 2009 VOL 150 ISS 6 FILE LAST UPDATED: 3 Feb 2009 (20090203/ED)

Caplus now includes complete International Patent Classification (IPC) reclassification data for the third quarter of 2008.

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This file contains CAS Registry Numbers for easy and accurate substance identification.

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L4 20 L3

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=> s 14 and calpain
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          6326 CALPAIN
                 (CALPAIN OR CALPAINS)
           16 L4 AND CALPAIN
L5
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    ANSWER 1 OF 10 CAPLUS COPYRIGHT 2009 ACS on STN
ACCESSION NUMBER:
                     2008:697982 CAPLUS
DOCUMENT NUMBER:
                        149:54264
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TITLE: Preparation of 2-hydroxytetrahydrofuran peptide

derivatives for use as medicaments

INVENTOR(S): Auvin, Serge; Chabrier de Lassauniere, Pierre-Etienne PATENT ASSIGNEE(S): Societe de Conseils de Recherches et d'Applications

Scientifiques (S.C.R.A.S.), Fr.

U.S., 20pp., Cont.-in-part of U.S. Ser. No. 532,731. CODEN: USXXAM SOURCE:

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

| PA | IENT | NO. | | | KIN | | DATE | | | APPL | ICAT | ION : | NO. | | D. | ATE | | |
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| | 7384 | | | | В2 | | | 0610 | | US 2 | 005- | 1154 | 80 | | 2 | 0050 | 427 < | |
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| | 2863 | | | | A1 | | | 0610 | | FR 2 | 003- | 1436 | 8 | | 2 | 0031 | 209 < | |
| | 2863 | | | | | | | 0224 | | | | | | | | | | |
| WO | 2005 | 0565 | 51 | | A2 | | 2005 | 0623 | | WO 2 | 004 - 1 | FR31 | 47 | | 2 | 0041 | 208 < | |
| WO | 2005 | 0565 | 51 | | A3 | | 2005 | 0811 | | | | | | | | | | |
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| | | CN, | CO, | CR, | CU, | CZ, | DE, | DK, | DM, | DZ, | EC, | EE, | EG, | ES, | FI, | GB, | GD, | |
| | | GE, | GH, | GM, | HR, | HU, | ID, | IL, | IN, | IS, | JP, | KE, | KG, | KP, | KR, | KZ, | LC, | |
| | | LK. | LR. | LS, | LT. | LU, | LV. | MA. | MD, | MG. | MK. | MN. | MW. | MX, | MZ, | NA. | NI. | |
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| | RW: | | | | | | | | | , | | | | | ZM, | | | |
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| | | | | | TD, | | DI, | ъо, | CL, | CO, | C1, | CI1, | OH, | OIV, | 02, | OW, | 1111, | |
| TTC | 2006 | | • | | | | 2006 | 0727 | | 110 2 | 005 | 5227 | 21 | | 2 | 0050 | 426 < | |
| | | | | | | | | | | 05 2 | 005- | 3327 | 31 | | | 0050 | 420 < | |
| | 7465 | | | | | | 2008 | 1710 | | o | 000 | 1 126 | ^ | | 3 0 | 0001 | 200 | |
| PRIORIT | Y APP | LN. | TNEO | .: | | | | | | | | | | | | | 209 < | |
| | | | | | | | | | | | | | | | | | 208 < | |
| | | | | | | | | | | US 2 | 005- | 5327 | 31 | | A2 2 | 0050 | 426 | |
| OTHER SO | OURCE | (S): | | | MARI | PAT | 149: | 5426 | 4 | | | | | | | | | |

OTHER SOURCE(S): MARPAT 149:54264

GΙ

$$R^{4}$$
 R^{3}
 R^{1}
 R^{5}
 R^{6}
 R^{2}
 R^{2}
 R^{1}
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 R^{3}
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 R^{4}
 R^{3}
 R^{4}
 R^{5}
 R^{5

AΒ The invention relates to hydroxytetrahydrofuran derivs. I [R1, R2, R4, R5, R6 are independently H, halo, OH, alkyl, alkoxy, cyano, nitro, amino or acylamino groups; R3 is H, alkyl, acyl, or carbalkoxy; W is a bond, CH2CH2, CH:CH, O, S, NH or alkylimino; X is CO, Y-CO, O-Y-CO (Y is alkylene or haloalkylene), NH, alkylimino, acyl, or carbalkoxy; AA is NR7(CH2)3CHR8CO (R7, R8 are H or alkyl), a natural amino acid, including a natural amino acid whose side chain carries a protected reactive chemical function; n is 2 or 3; R is H, alkyl, or alkanoyl] which have calpain-inhibiting activity and/or activity which traps reactive oxygen species and are useful for treating inflammatory and immunol. diseases, cardiovascular and cerebrovascular diseases, disorders of the central or peripheral nervous system, osteoporosis, muscular dystrophy, proliferative diseases, cataract, rejection reactions following organ transplants and autoimmune and viral diseases. Thus, N-(10H-phenothiazin-2-ylcarbonyl)-L-leucyl-L-leucyl-N1-[(3S)-2hydroxytetrahydrofuran-3-yl]-L-leucinamide, prepared by a multistep sequence which starts with reaction of Cbz-protected L-leucine with (S)-2-amino-4-butyrolactone hydrobromide, showed IC50 \leq 5 μM in the human calpain I inhibition assay.

IT 853208-13-6P

CN

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of hydroxytetrahydrofuran peptide derivs. for use as medicaments)

RN 853208-13-6 CAPLUS

L-Leucinamide, N-(10H-phenothiazin-2-ylcarbonyl)-L-leucyl-N-[(3S)-2-(acetyloxy)tetrahydro-3-furanyl]- (CA INDEX NAME)

Absolute stereochemistry.

REFERENCE COUNT: 6 THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 2 OF 10 CAPLUS COPYRIGHT 2009 ACS on STN

ACCESSION NUMBER: 2005:1075636 CAPLUS

DOCUMENT NUMBER: 143:339689

TITLE: Use of a phenothiazine derivative for preventing

and/or treating hearing loss

INVENTOR(S): Pignol, Bernadette; Puel, Jean-Luc; Auvin, Serge;

Chabrier de Lassauniere, Pierre-Etienne; Wang, Jing

PATENT ASSIGNEE(S): Societe de Conseils de Recherches et d'Applications

Scientifiques S.C.R.A.S., Fr.

SOURCE: PCT Int. Appl., 33 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: French

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

| PAT | PATENT NO. | | | | | | KIND DATE | | | APPLICATION NO. | | | | | | | | | | | |
|---------|------------|------|------|-----|------------|-----|-----------|------|-----|-----------------|------|------|-----|-----|-----|------|-----|----|--|--|--|
| WO | 2005 | 0923 | 45 | | A1 | | | | | | | | | | | 0050 | 325 | < | | | |
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| | | CN, | CO, | CR, | CU, | CZ, | DE, | DK, | DM, | DZ, | EC, | EE, | EG, | ES, | FΙ, | GB, | GD, | | | | |
| | | GE, | GH, | GM, | HR, | HU, | ID, | IL, | IN, | IS, | JP, | KE, | KG, | KP, | KR, | KZ, | LC, | | | | |
| | | LK, | LR, | LS, | LT, | LU, | LV, | MA, | MD, | MG, | MK, | MN, | MW, | MX, | MZ, | NA, | ΝI, | | | | |
| | | NO, | NZ, | OM, | PG, | PH, | PL, | PT, | RO, | RU, | SC, | SD, | SE, | SG, | SK, | SL, | SM, | | | | |
| | | SY, | ТJ, | TM, | TN, | TR, | TT, | TZ, | UA, | UG, | US, | UΖ, | VC, | VN, | YU, | ZA, | ZM, | ZW | | | |
| | RW: | BW, | GH, | GM, | ΚE, | LS, | MW, | MZ, | NΑ, | SD, | SL, | SZ, | TZ, | UG, | ZM, | ZW, | ΑM, | | | | |
| | | ΑZ, | BY, | KG, | ΚZ, | MD, | RU, | ТJ, | TM, | AT, | ΒE, | BG, | CH, | CY, | CZ, | DE, | DK, | | | | |
| | | EE, | ES, | FI, | FR, | GB, | GR, | HU, | ΙE, | IS, | ΙΤ, | LT, | LU, | MC, | NL, | PL, | PT, | | | | |
| | | RO, | SE, | SI, | SK, | TR, | BF, | ВJ, | CF, | CG, | CI, | CM, | GA, | GN, | GQ, | GW, | ML, | | | | |
| | | MR, | NE, | SN, | TD, | ΤG | | | | | | | | | | | | | | | |
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| FR | 2867 | 979 | | | В1 | | 2006 | 0630 | | | | | | | | | | | | | |
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| EP | 1732 | | | | В1 | | 2008 | | | | | | | | | | | | | | |
| | R: | ΑT, | ΒE, | BG, | CH, | CY, | CZ, | DE, | DK, | ΕE, | ES, | FΙ, | FR, | GB, | GR, | ΗU, | ΙE, | | | | |
| | | IS, | ΙΤ, | LI, | LT, | LU, | MC, | ΝL, | PL, | PT, | RO, | SE, | SI, | SK, | TR | | | | | | |
| | 1933 | | | | | | 2007 | | | | | | | | | | | | | | |
| | 2007 | | | | | | | | | | | | | | 2 | | | | | | |
| US | 2008 | 0275 | 034 | | A1 | | 2008 | 1106 | | | | | | | | | | | | | |
| PRIORIT | Y APP | LN. | INFO | .: | | | | | | FR 2 | 004- | 3203 | | 2 | A 2 | 0040 | 329 | < | | | |
| | | | | | | | | | | | | | | | A 2 | 0040 | 614 | < | | | |
| | | | | | | | | | | WO 2 | 005- | FR71 | 3 | Ī | W 2 | 0050 | 325 | | | | |

OTHER SOURCE(S): MARPAT 143:339689

GI

AB The invention discloses the use of a phenothiazine derivative I (R = H, alkyl, aralkyl, etc.) for preparing a medicine for preventing and/or treating hearing loss.

IT 742104-24-1

RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

Ι

(phenothiazine derivs. for prevention and/or treatment of hearing loss)

RN 742104-24-1 CAPLUS

CN 10H-Phenothiazine-2-carboxamide, N-[(1S)-1-[[[(3S)-2-(acetyloxy)tetrahydro-3-furanyl]amino]carbonyl]-3-methylbutyl]- (CA INDEX NAME)

REFERENCE COUNT: 10 THERE ARE 10 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 3 OF 10 CAPLUS COPYRIGHT 2009 ACS on STN

ACCESSION NUMBER: 2005:492131 CAPLUS

DOCUMENT NUMBER: 143:44075

TITLE: Preparation of peptidyl 3-aminotetrahydro-2-furanol

derivatives for use as drugs

INVENTOR(S): Auvin, Serge; Chabrier De Lassauniere, Pierre Etienne

PATENT ASSIGNEE(S): Societe De Conseils de Recherches et d'Applications

Scientifiques SCRAS, Fr.

SOURCE: Fr. Demande, 53 pp.

CODEN: FRXXBL

DOCUMENT TYPE: Patent LANGUAGE: French

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

| PA | TENT | NO. | | | KIN | D | DATE | | | | ICAT | | | | D | ATE | | |
|----------|--------------|-------|------|-----|-------|-------|--------------|------|-----|------|------|------|-----|-----|------|------|-------|--|
| | 2863 2863 | | | | | | 2005 2006 | | | | | | | | 2 | 0031 | 209 < | |
| | 2548 | | | | | | | | | CA 2 | 004- | 2548 | 448 | | 2 | 0041 | 208 < | |
| WO | 2005 | 0565 | 51 | | A2 | | 2005 | 0623 | | WO 2 | 004- | FR31 | 47 | | 2 | 0041 | 208 < | |
| | 2005 | | | | | | | | | | | | | | | | | |
| _ | | | | | | | AU, | | | BB, | BG, | BR, | BW. | BY, | BZ, | CA, | CH, | |
| | | | | | | | DE, | | | | | | | | - | - | | |
| | | | | | | | ID, | | | | | | | | | | | |
| | | | | | | | LV, | | | | | | | | | | | |
| | | | | | | | PL, | | | | | | | | | | | |
| | | ΤJ, | TM, | TN, | TR, | TT, | TZ, | UA, | UG, | US, | UZ, | VC, | VN, | YU, | ZA, | ZM, | ZW | |
| | RW: | BW, | GH, | GM, | KE, | LS, | MW, | MZ, | NA, | SD, | SL, | SZ, | TZ, | UG, | ZM, | ZW, | AM, | |
| | | | | | | | RU, | | | | | | | | | | | |
| | | EE, | ES, | FΙ, | FR, | GB, | GR, | ΗU, | ΙE, | IS, | ΙΤ, | LT, | LU, | MC, | NL, | PL, | PT, | |
| | | RO, | SE, | SI, | SK, | TR, | BF, | ВJ, | CF, | CG, | CI, | CM, | GΑ, | GN, | GQ, | GW, | ML, | |
| | | MR, | ΝE, | SN, | TD, | | | | | | | | | | | | | |
| EP | 1701 | 974 | | | A2 | | 2006 | 0920 | | EP 2 | 004- | 8163 | 63 | | 2 | 0041 | 208 < | |
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| | R: | , | , | | | , | | , | | | , | | , | | | MC, | PT, | |
| | | | | | | | CY, | | | | | | | | | | | |
| JP | 2007 | 5139 | 29 | | Τ | | 2007 | 0531 | | JP 2 | 006- | 5435 | 81 | | 2 | 0041 | 208 < | |
| | | | | | | | | | | | | | | | | | 208 < | |
| | | | | | | | | | | | | | | | | | 208 < | |
| | 2006 | | | | | | | | | US 2 | 005- | 5327 | 31 | | 2 | 0050 | 426 < | |
| | 7465 | | | | | | 2008 | | | | | | | | | | | |
| | 7384 | | | | | | 2008 | | | US 2 | 005- | 1154 | 80 | | 2 | 0050 | 427 < | |
| | 2005 | | | | A1 | | 2005 | 1006 | | | | | _ | | _ | | | |
| PRIORIT | Y APP | LN. | INFO | .: | | | | | | | | | | | | | 209 < | |
| | | | | | | | | | | - | | _ | | | | | 208 < | |
| OTHER SO | OLIDOE | (C) : | | | MENTO | ייי ע | 1/2: | 4407 | | US 2 | 005- | 5327 | 3 I | | A2 2 | 0050 | 426 | |

OTHER SOURCE(S): MARPAT 143:44075

GΙ

$$A-X-(AA)_{n}^{-N}$$

AB The invention relates to peptide derivs. I [A is (un)substituted carbazolyl, dibenzo[b,f]azepinyl or 10,11-dihydro derivs., phenoxazinyl, phenothiazinyl or phenazinyl; X is CO, Y-CO, O-Y-CO or NR1-Y-CO; Y is alkylene or haloalkylene; R, R1 are independently H, alkyl or acyl; AA is a natural amino acid or derivative; n is 2,3] or their salts which inhibit calpains and lipid peroxidn. and can be used to treat inflammatory, immunol., cardiovascular and other diseases. Thus, N-(10H-phenothiazin-2-ylcarbonyl)-L-leucyl-L-leucyl-N1-[(3S)-2-hydroxytetrahydrofuran-3-yl]-L-leucinamide was prepared by a multistep procedure involving reactions of Cbz-L-Leucine (Cbz = benzyloxycarbonyl), (S)-2-amino-4-butyrolactone hydrobromide, and 2-acetylphenothiazine and treatment with 2N HCl. The product showed IC50 < 5 μM for inhibition of human calpain I.

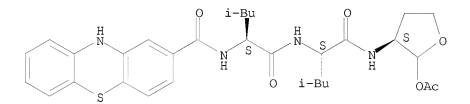
IT 853208-13-6P

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of peptidyl aminotetrahydrofuranol derivs. for use as drugs) ${\tt RN} - 853208-13-6 - {\tt CAPLUS}$

CN L-Leucinamide, N-(10H-phenothiazin-2-ylcarbonyl)-L-leucyl-N-[(3S)-2-(acetyloxy)tetrahydro-3-furanyl]- (CA INDEX NAME)

Absolute stereochemistry.



REFERENCE COUNT: 1 THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 4 OF 10 CAPLUS COPYRIGHT 2009 ACS on STN

ACCESSION NUMBER: 2004:498592 CAPLUS

DOCUMENT NUMBER: 141:207514

TITLE: Novel dual inhibitors of calpain and lipid

peroxidation

AUTHOR(S): Auvin, Serge; Pignol, Bernadette; Navet, Edith; Pons,

Dominique; Marin, Jean-G.; Bigg, Dennis; Chabrier,

Pierre-E.

CORPORATE SOURCE: Department of Medicinal Chemistry, Ipsen Research

Laboratories, Institut Henri Beaufour, Les Ulis,

91966, Fr.

SOURCE: Bioorganic & Medicinal Chemistry Letters (2004

), 14(14), 3825-3828

CODEN: BMCLE8; ISSN: 0960-894X

PUBLISHER: Elsevier Science B.V.

DOCUMENT TYPE: Journal

LANGUAGE: English

OTHER SOURCE(S): CASREACT 141:207514

GI

AB A series of mols. I (R1 = phenothiazin-1-yl, phenothiazin-2-yl, 1-benzyl-5-indolinylamino, etc., R2 = H; R1 = phenothiazin-2-yl, R2 = MeCO) with dual inhibitory activities on calpain and lipid peroxidn. were synthesized. These hybrid compds. were built on the calpain pharmacophore 2-hydroxytetrahydrofuran linked to a set of antioxidants via a L-leucine linker. I (R1 = phenothiazin-2-yl, R2 = MeCO), the most potent in cellular calpain and lipid peroxidn. inhibitions, provided effective protection against glial cell death induced by maitotoxin.

IT 742104-24-1P, BN 82270

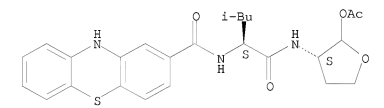
RL: PAC (Pharmacological activity); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)

(preparation of (peptidyl)(hydroxy)tetrahydrofurans as dual inhibitors of calpain and lipid peroxidn.)

RN 742104-24-1 CAPLUS

CN 10H-Phenothiazine-2-carboxamide, N-[(1S)-1-[[[(3S)-2-(acetyloxy)tetrahydro-3-furanyl]amino]carbonyl]-3-methylbutyl]- (CA INDEX NAME)

Absolute stereochemistry.



REFERENCE COUNT: 15 THERE ARE 15 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 5 OF 10 CAPLUS COPYRIGHT 2009 ACS on STN

ACCESSION NUMBER: 2001:338525 CAPLUS

DOCUMENT NUMBER: 134:353248

TITLE: Novel heterocyclic compounds and their use as

medicines

INVENTOR(S): Auvin, Serge; Chabrier De Lassauniere, Pierre-Etienne PATENT ASSIGNEE(S): Societe De Conseils De Recherches Et D'applications

Scientifiques (S.C.R.A.S.), Fr.

SOURCE: PCT Int. Appl., 77 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: French

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

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WO 2001032654 A2 20010510 WO 2001032654 A3 20010927
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            HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT,
            LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU,
            SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN,
            YU, ZA, ZW
        RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY,
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                       MARPAT 134:353248
OTHER SOURCE(S):
    Novel heterocyclic derivs. which have calpain inhibiting and/or
AB
    reactive oxygen species trapping activity (no data) are reported. Thus,
     (R)-Trolox was treated with (S)-2-aminobutyrolactone hydrochloride,
    followed by DIBAL reduction to give (2R)-6-hydroxy-N-[(3S)-2-
    hydroxytetrahydrofuran-3-y1]-2,5,7,8-tetramethyl-3,4-dihydro-2H-chromene-2-
    carboxamide.
    339007-48-6P 339007-52-2P 339007-53-3P
IT
    339007-55-5P 339007-56-6P 339007-76-0P
    RL: SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological
    study); PREP (Preparation); USES (Uses)
        (preparation of novel heterocyclic compds. as calpain inhibitors
       and trapping agents for reactive oxygen species)
RN
    339007-48-6 CAPLUS
CN
    10H-Phenothiazine-2-carboxamide, N-[(1S)-1-[[(2S,3S)-2-(2S,3S)-1]]]
```

(acetyloxy)tetrahydro-3-furanyl]amino]carbonyl]-3-methylbutyl]- (CA INDEX NAME)

Absolute stereochemistry.

RN 339007-52-2 CAPLUS

CN Propanoic acid, 2,2-dimethyl-, (3S)-tetrahydro-3-[[(2S)-4-methyl-1-oxo-2-[(10H-phenothiazin-2-ylcarbonyl)amino]pentyl]amino]-2-furanyl ester (CA INDEX NAME)

Absolute stereochemistry.

RN 339007-53-3 CAPLUS

CN Butanoic acid, 3,3-dimethyl-, (3S)-tetrahydro-3-[[(2S)-4-methyl-1-oxo-2-[(10H-phenothiazin-2-ylcarbonyl)amino]pentyl]amino]-2-furanyl ester (CA INDEX NAME)

Absolute stereochemistry.

RN 339007-55-5 CAPLUS

CN Benzeneacetic acid, (3S)-tetrahydro-3-[[(2S)-4-methyl-1-oxo-2-[(10H-phenothiazin-2-ylcarbonyl)amino]pentyl]amino]-2-furanyl ester (CA INDEX NAME)

RN 339007-56-6 CAPLUS

CN L-Phenylalanine, N,N-dimethyl-, (3S)-tetrahydro-3-[[(2S)-4-methyl-1-oxo-2-[(10H-phenothiazin-2-ylcarbonyl)amino]pentyl]amino]-2-furanyl ester (CA INDEX NAME)

Absolute stereochemistry.

RN 339007-76-0 CAPLUS

CN 10H-Phenothiazine-2-carboxamide, N-[(1S)-1-[[((2R,3S)-2-(acetyloxy)tetrahydro-3-furanyl]amino]carbonyl]-3-methylbutyl]- (CA INDEX NAME)

Absolute stereochemistry.

REFERENCE COUNT: 6 THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 6 OF 10 CAPLUS COPYRIGHT 2009 ACS on STN

ACCESSION NUMBER: 1999:233912 CAPLUS

DOCUMENT NUMBER: 130:252373

TITLE: Preparation and formulation of O-containing

heterocyclic derivatives as cysteine protease

inhibitors

INVENTOR(S): Usui, Yoshihiro; Masuda, Hirokazu; Ando, Naoko; Nakao,

Akira; Ando, Ryoichi; Yoshii, Narihiko; Saito,

Ken-ichi

PATENT ASSIGNEE(S): Mitsubishi Chemical Corporation, Japan

SOURCE: PCT Int. Appl., 115 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

GΙ

| PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|------------------------|--------|--------------|-------------------|-----------------|
| | | | | |
| WO 9916761 | A1 | 19990408 | WO 1998-JP4420 | 19980930 < |
| W: CA, CN, KF | US | | | |
| RW: AT, BE, CH | CY, DE | , DK, ES, FI | , FR, GB, GR, IE, | IT, LU, MC, NL, |
| PT, SE | | | | |
| JP 11171881 | A | 19990629 | JP 1998-277586 | 19980930 < |
| PRIORITY APPLN. INFO.: | | | JP 1997-266034 | A 19970930 < |
| OTHER SOURCE(S): | MARPAT | 130:252373 | | |
| CT | | | | |

AΒ The title compds. I [R1 represents optionally substituted C6-14 aryl or an optionally substituted heterocycle residue; R2 represents hydrogen or C1-10 alkyl optionally substituted by C6-14 aryl; R3 represents hydrogen or R4CO (R4 represents C1-10 alkyl); and A represents C1-3 alkylene optionally substituted by C1-3 alkyl] are prepared I are useful as cysteine protease inhibitors excellent in oral absorbability, migration to tissues, and can easily pass through the cell membrane, etc. (3S)-3-[(S)-2-(4,6-dimethoxy-2-pyrimidiny1) amino-4-methylvalerylamino]-2tetrahydrofuranol in vitro showed IC50 of 1.27 μM against calpain.

ΙT 221683-02-9P 221683-09-6P 221683-12-1P 221683-20-1P

> RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of O-containing heterocyclic derivs. as cysteine protease inhibitors)

221683-02-9 CAPLUS RN

Pentanamide, N-[(3S)-2-(acetyloxy)tetrahydro-3-furanyl]-2-[(4,6-dimethoxy-CN 2-pyrimidinyl)amino]-4-methyl-, (2S)- (CA INDEX NAME)

CN Pentanamide, N-[(3S)-2-(acetyloxy)tetrahydro-3-furanyl]-4-methyl-2-[[2-nitro-4-(trifluoromethyl)phenyl]amino]-, (2S)- (CA INDEX NAME)

Absolute stereochemistry.

RN 221683-12-1 CAPLUS

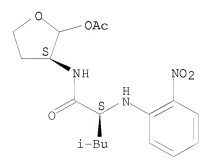
CN Pentanamide, N-[(3S)-2-(acetyloxy)tetrahydro-3-furanyl]-2-(2-benzoxazolylamino)-4-methyl-, (2S)- (CA INDEX NAME)

Absolute stereochemistry.

RN 221683-20-1 CAPLUS

CN Pentanamide, N-[(3S)-2-(acetyloxy)tetrahydro-3-furanyl]-4-methyl-2-[(2-nitrophenyl)amino]-, (2S)- (CA INDEX NAME)

Absolute stereochemistry.



REFERENCE COUNT: 6 THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 7 OF 10 CAPLUS COPYRIGHT 2009 ACS on STN

ACCESSION NUMBER: 1998:246624 CAPLUS

DOCUMENT NUMBER: 129:32318

ORIGINAL REFERENCE NO.: 129:6761a,6764a

TITLE: Cataract curative medicine.

INVENTOR(S): Watanabe, Toshiaki; Yoshii, Shigehiko; Saito, Kenichi;

Ando, Ryoichi

PATENT ASSIGNEE(S): Mitsubishi Chemical Industries Ltd., Japan

SOURCE: Jpn. Kokai Tokkyo Koho, 74 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

| | PATENT NO. | KIND | DATE | APPLICATION NO. | | DATE | | | | | |
|------------------|--|--------------|-------------------------|--|--|--------------------------|--|--|--|--|--|
| | DRITY APPLN. INFO.: ER SOURCE(S): | A MARPA] | 19980421 T 129:32318 | JP 1997-197216 JP 1996-208540 | | 19970723 < 19960807 < | | | | | |
| AB | For diagram(s), see printed CA Issue. The cataract curative medicine has an effective component of structure (I), its salt, solvate, or hydrate, where R1 is R4-C0-, R4-O-C0-, or R4-S02- (R4: C1-20 alkyl), R2 is C1-C6 alkyl, R3 is H or R5-C0- (R5: C1-10 alkyl), and A is C1-3 alkylene. Thus, 998 mg N-phenylsulfonyl-L-leucine was react with 6 mL S02C12 and 443 mg homoserine lactone to give (S)-3-[(S)-4-methyl-2-phenylsulfonylaminovalerylamino]-2-tetrahydrofuranone 861 mg, which was reacted with hydrogendiisobutylaluminum to give (3S)-3-[(S)-4-Methyl-2-phenylsulfonylaminovalerylamino]-2-tetrahydrofuranol 191 mg, which showed strong calpain inhibition | | | | | | | | | | |
| IT | activity (I C50 0.62 μM). 167765-43-7P 201155-39-7DP, salts, solvates, or hydrates 201155-39-7P 201155-40-0DP, salts, solvates, or hydrates 201155-40-0P 201155-41-1DP, salts, solvates, or hydrates 201155-41-1P 201155-42-2P 201155-44-4DP, salts, solvates, or hydrates 201155-44-4P 201155-46-6P 201155-47-7DP, salts, solvates, or hydrates 201155-47-7P 201155-49-9P 201155-50-2P 201155-52-4DP, salts, solvates, or hydrates 201155-52-4P 201155-54-6DP , salts, solvates, or hydrates 201155-54-6DP 201155-58-0DP, salts, solvates, or hydrates 201155-58-0P 201155-60-4DP, salts, solvates, or hydrates 201155-60-4P 201157-12-2DP, salts, solvates, or hydrates 201157-12-2P 201157-68-8P 207500-74-1P 207500-75-2P 207500-76-3DP, salts, solvates, or hydrates 207500-76-3P RL: BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) | | | | | | | | | | |
| R N CN | | S S)-1-[[| [(2S,3S)-2-(a | acetyloxy)tetrahydro- -, phenylmethyl este | | 9CI) (CA | | | | | |

RN 201155-39-7 CAPLUS

CN Carbamic acid, [(1S)-1-[[[(2S,3S)-2-(acetyloxy)tetrahydro-3-furanyl]amino]carbonyl]-3-methylbutyl]-, tetradecyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 201155-39-7 CAPLUS

CN Carbamic acid, [(1S)-1-[[[(2S,3S)-2-(acetyloxy)tetrahydro-3-furanyl]amino]carbonyl]-3-methylbutyl]-, tetradecyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 201155-40-0 CAPLUS

CN Pentadecanamide, N-[(1S)-1-[[[(2S,3S)-2-(acetyloxy)tetrahydro-3-(acetylox

furanyl]amino]carbonyl]-3-methylbutyl]- (CA INDEX NAME)

Absolute stereochemistry.

RN 201155-40-0 CAPLUS

CN Pentadecanamide, N-[(1S)-1-[[[(2S,3S)-2-(acetyloxy)tetrahydro-3-furanyl]amino]carbonyl]-3-methylbutyl]- (CA INDEX NAME)

Absolute stereochemistry.

RN 201155-41-1 CAPLUS

CN Hexadecanamide, N-[(1S)-1-[[[(2S,3S)-2-(acetyloxy)tetrahydro-3-furanyl]amino]carbonyl]-3-methylbutyl]- (CA INDEX NAME)

Absolute stereochemistry.

RN 201155-41-1 CAPLUS

CN Hexadecanamide, N-[(1S)-1-[[[(2S,3S)-2-(acetyloxy)tetrahydro-3-furanyl]amino]carbonyl]-3-methylbutyl]- (CA INDEX NAME)

Absolute stereochemistry.

RN 201155-42-2 CAPLUS

CN Benzamide, N-[(1S)-1-[[[(2S,3S)-2-(acetyloxy)tetrahydro-3-furanyl]amino]carbonyl]-3-methylbutyl]-2-fluoro- (CA INDEX NAME)

Absolute stereochemistry.

RN 201155-44-4 CAPLUS

CN Pentanamide, N-[(2S,3S)-2-(acetyloxy)tetrahydro-3-furanyl]-2-[(hexadecylsulfonyl)amino]-4-methyl-, (2S)- (CA INDEX NAME)

Absolute stereochemistry.

RN 201155-44-4 CAPLUS

CN Pentanamide, N-[(2S,3S)-2-(acetyloxy)tetrahydro-3-furanyl]-2-[(hexadecylsulfonyl)amino]-4-methyl-, (2S)- (CA INDEX NAME) Absolute stereochemistry.

RN 201155-46-6 CAPLUS

CN Benzamide, N-[(1S)-1-[[[(2S,3S)-2-(acetyloxy)tetrahydro-3-furanyl]amino]carbonyl]-3-methylbutyl]-4-chloro- (CA INDEX NAME)

Absolute stereochemistry.

RN 201155-47-7 CAPLUS

CN Pentanamide, N-[(2S,3S)-2-(acetyloxy)tetrahydro-3-furanyl]-4-methyl-2-[[(2,4,6-trimethylphenyl)sulfonyl]amino]-, (2S)- (CA INDEX NAME)

Absolute stereochemistry.

RN 201155-47-7 CAPLUS

CN Pentanamide, N-[(2S,3S)-2-(acetyloxy)tetrahydro-3-furanyl]-4-methyl-2-[[(2,4,6-trimethylphenyl)sulfonyl]amino]-, (2S)- (CA INDEX NAME)

RN 201155-49-9 CAPLUS

CN Pentanamide, N-[(2S,3S)-2-(acetyloxy)tetrahydro-3-furanyl]-2-[[(4-methoxyphenyl)sulfonyl]amino]-4-methyl-, (2S)- (CA INDEX NAME)

Absolute stereochemistry.

RN 201155-50-2 CAPLUS

CN Pentanamide, N-[(2S,3S)-2-(acetyloxy)tetrahydro-3-furanyl]-4-methyl-2-[(2-naphthalenylsulfonyl)amino]-, (2S)- (CA INDEX NAME)

Absolute stereochemistry.

RN 201155-52-4 CAPLUS

CN Heptadecanamide, N-[(1S)-1-[[[(2S,3S)-2-(acetyloxy)tetrahydro-3-furanyl]amino]carbonyl]-3-methylbutyl]- (CA INDEX NAME)

RN 201155-52-4 CAPLUS

CN Heptadecanamide, N-[(1S)-1-[[[(2S,3S)-2-(acetyloxy)tetrahydro-3-furanyl]amino]carbonyl]-3-methylbutyl]- (CA INDEX NAME)

Absolute stereochemistry.

RN 201155-54-6 CAPLUS

CN Carbamic acid, [(1S)-1-[[[(2S,3S)-2-(acetyloxy)tetrahydro-3-furanyl]amino]carbonyl]-3-methylbutyl]-, tridecyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 201155-54-6 CAPLUS

CN Carbamic acid, [(1S)-1-[[[(2S,3S)-2-(acetyloxy)tetrahydro-3-furanyl]amino]carbonyl]-3-methylbutyl]-, tridecyl ester (9CI) (CA INDEX

NAME)

Absolute stereochemistry.

RN 201155-58-0 CAPLUS

CN Pentanamide, N-[(2S,3S)-2-(acetyloxy)tetrahydro-3-furanyl]-4-methyl-2-[(tetradecylsulfonyl)amino]-, (2S)- (CA INDEX NAME)

Absolute stereochemistry.

RN 201155-58-0 CAPLUS

CN Pentanamide, N-[(2S,3S)-2-(acetyloxy)tetrahydro-3-furanyl]-4-methyl-2-[(tetradecylsulfonyl)amino]-, (2S)- (CA INDEX NAME)

Absolute stereochemistry.

RN 201155-60-4 CAPLUS

CN Pentanamide, N-[(2S,3S)-2-(acetyloxy)tetrahydro-3-furanyl]-4-methyl-2-[(pentadecylsulfonyl)amino]-, (2S)- (CA INDEX NAME)

Absolute stereochemistry.

RN 201155-60-4 CAPLUS

CN Pentanamide, N-[(2S,3S)-2-(acetyloxy)tetrahydro-3-furanyl]-4-methyl-2-[(pentadecylsulfonyl)amino]-, (2S)- (CA INDEX NAME)

Absolute stereochemistry.

RN 201157-12-2 CAPLUS

CN Pentanamide, N-[(2S,3S)-2-(acetyloxy)tetrahydro-3-furanyl]-4-methyl-2-[(phenylsulfonyl)amino]-, (2S)- (CA INDEX NAME)

RN 201157-12-2 CAPLUS

Pentanamide, N-[(2S,3S)-2-(acetyloxy)tetrahydro-3-furany1]-4-methyl-2-CN [(phenylsulfonyl)amino]-, (2S)- (CA INDEX NAME)

Absolute stereochemistry.

RN

201157-68-8 CAPLUS Pentanamide, N-[(2S,3S)-2-(acetyloxy)tetrahydro-3-furany1]-2-[[(4- $^{\circ}$ CN chlorophenyl)sulfonyl]amino]-4-methyl-, (2S)- (CA INDEX NAME)

Absolute stereochemistry.

RN 207500-74-1 CAPLUS

Pentanamide, 4-methyl-2-[(phenylsulfonyl)amino]-N-[(2S,3S)-tetrahydro-2-(1-CN oxopropoxy)-3-furany1]-, (2S)- (CA INDEX NAME)

Absolute stereochemistry.

207500-75-2 CAPLUS RN

Pentanamide, 2-[[[4-(1,1-dimethylethyl)phenyl]sulfonyl]amino]-4-methyl-N-CN [(2S,3S)-tetrahydro-2-(1-oxopropoxy)-3-furanyl]-, (2S)- (CA INDEX NAME)

Absolute stereochemistry.

RN 207500-76-3 CAPLUS

CN Carbamic acid, [(1S)-1-[[[(2S,3S)-2-(acetyloxy)tetrahydro-3-furanyl]amino]carbonyl]-3-methylbutyl]-, heptadecyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 207500-76-3 CAPLUS

CN Carbamic acid, [(1S)-1-[[[(2S,3S)-2-(acetyloxy)tetrahydro-3-furanyl]amino]carbonyl]-3-methylbutyl]-, heptadecyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L6 ANSWER 8 OF 10 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1998:102857 CAPLUS

DOCUMENT NUMBER: 128:167712

ORIGINAL REFERENCE NO.: 128:33065a,33068a

TITLE: Preparation of oxygenic heterocyclic derivatives of

amino acid amides as cysteine protease inhibitors

INVENTOR(S): Ando, Ryoichi; Masuda, Hirokazu; Aritomo, Keiichi;

Yoshii, Narihiko; Saito, Ken-Ichi

PATENT ASSIGNEE(S): Mitsubishi Chemical Corporation, Japan

SOURCE: PCT Int. Appl., 68 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

W: CA, CN, JP, KR, US

RW: AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE

PRIORITY APPLN. INFO.: JP 1996-199037 A 19960729 <--

OTHER SOURCE(S): MARPAT 128:167712

GΙ

AB Oxygenic heterocyclic derivs. of general formula [I; R1 = R4CO, R4O2C, R4SO2 (R4 = straight-chain C11-20 alkyl); R2 = C1-10 alkyl optionally substituted by C6-14 aryl; R3 = H, R5CO (wherein R5 = C1-10 alkyl); A = C1-3 alkylene optionally substituted by C1-3 alkyl], salts thereof, and solvates or hydrates thereof are prepared These compds. exhibit a potent inhibitory activity against cysteine proteases such as calpain, papain, cathepsin B, cathepsin H, cathepsin L, calpain, and interleukin 1β -converting enzyme and are excellent in absorbability through oral administration, tissue transportability, and cell membrane permeability and are useful for the treatment of muscular dystrophy, muscular atrophy, myocardial infarction, stroke, Alzheimer's disease, disorders of cognition and motor disorders in head trauma, multiple sclerosis, neuropathy of peripheral nerve, cataract, allergy, hepatitis siderans, osteoporosis, hypercalcemia, breast cancer, prostate cancer, prostatomegaly, inhibitors of cancer proliferation and metastasis, and blood platelet aggregation inhibitors. Thus, (3S)-3-[(S)-2-(tert-butoxycarbonylamino)-4-methylvalerylamino]-2tetrahydrofuranone was stirred with 4 N HCl in EtOAc at room temperature for 45 min and then acylated by heptadecanoyl chloride in the presence of Et3N in CH2Cl2 at room temperature overnight to give (3S)-3-[(S)-2-(heptadecanoylamino)-4-methylvalerylamino]-2tetrahydrofuranone, which was reduced by LiAlH4 in THF at $-68\,^{\circ}$ for 1 h to give (3S)-[(N-heptadecanoyl-L-leucinyl)amino]-2-tetrahydrofuranol (II; R = heptadecanoyl). The latter compound and II (R =pentadecylsulfonyl) in vitro showed IC50 of 1.05 and 0.09 μM , resp., against m-calpain.

IT 201155-39-7P 201155-40-0P 201155-41-1P

Absolute stereochemistry.

RN 201155-40-0 CAPLUS
CN Pentadecanamide, N-[(1S)-1-[[[(2S,3S)-2-(acetyloxy)tetrahydro-3-furanyl]amino]carbonyl]-3-methylbutyl]- (CA INDEX NAME)

Absolute stereochemistry.

RN 201155-41-1 CAPLUS
CN Hexadecanamide, N-[(1S)-1-[[[(2S,3S)-2-(acetyloxy)tetrahydro-3-furanyl]amino]carbonyl]-3-methylbutyl]- (CA INDEX NAME)

RN 201155-44-4 CAPLUS

CN Pentanamide, N-[(2S,3S)-2-(acetyloxy)tetrahydro-3-furanyl]-2-[(hexadecylsulfonyl)amino]-4-methyl-, (2S)- (CA INDEX NAME)

Absolute stereochemistry.

RN 201155-52-4 CAPLUS

CN Heptadecanamide, N-[(1S)-1-[[[(2S,3S)-2-(acetyloxy)tetrahydro-3-furanyl]amino]carbonyl]-3-methylbutyl]- (CA INDEX NAME)

Absolute stereochemistry.

RN 201155-54-6 CAPLUS

CN Carbamic acid, [(1S)-1-[[[(2S,3S)-2-(acetyloxy)tetrahydro-3-furanyl]amino]carbonyl]-3-methylbutyl]-, tridecyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 201155-56-8 CAPLUS

CN Carbamic acid, [1-[[[2-(acetyloxy)tetrahydro-3-furanyl]amino]carbonyl]-3-methylbutyl]-, pentadecyl ester, [2S-[2 α ,3 α (R*)]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 201155-58-0 CAPLUS

CN Pentanamide, N-[(2S,3S)-2-(acetyloxy)tetrahydro-3-furanyl]-4-methyl-2-[(tetradecylsulfonyl)amino]-, (2S)- (CA INDEX NAME)

Absolute stereochemistry.

RN 201155-60-4 CAPLUS

CN Pentanamide, N-[(2S,3S)-2-(acetyloxy)tetrahydro-3-furanyl]-4-methyl-2-[(pentadecylsulfonyl)amino]-, (2S)- (CA INDEX NAME)

Absolute stereochemistry.

RN 202814-98-0 CAPLUS

CN Dodecanamide, N-[(1S)-1-[[[(2S,3S)-2-(acetyloxy)tetrahydro-3-furanyl]amino]carbonyl]-3-methylbutyl]- (CA INDEX NAME)

Absolute stereochemistry.

RN 202815-01-8 CAPLUS

CN Octadecanamide, N-[(1S)-1-[[[(2S,3S)-2-(acetyloxy)tetrahydro-3-furanyl]amino]carbonyl]-3-methylbutyl]- (CA INDEX NAME)

REFERENCE COUNT: 5 THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 9 OF 10 CAPLUS COPYRIGHT 2009 ACS on STN

ACCESSION NUMBER: 1998:65808 CAPLUS

DOCUMENT NUMBER: 128:102004

ORIGINAL REFERENCE NO.: 128:19985a,19988a

TITLE: Preparation of hydroxytetrahydrofuran derivatives as

remedies for ischemic diseases

INVENTOR(S): Yoshii, Narihiko; Saito, Ken-ichi; Kawasumi, Hisashi;

Anabuki, Jun; Ando, Ryoichi

PATENT ASSIGNEE(S): Mitsubishi Chemical Corp., Japan

SOURCE: PCT Int. Appl., 117 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

| PATENT NO. | | KIND DATE | | APPLICATION NO. | DATE 19970709 < | | | |
|-------------------------------------|----------------|-----------|----------------------|---|--|--|--|--|
| WO 9801130 W: US | | | 19980115 | WO 1997-JP2378 | | | | |
| RW: AT, JP 10101558 EP 925786 | | A A1 | 19980421 19990630 | FR, GB, GR, IE, IT, JP 1997-179756 EP 1997-930735 | LU, MC, NL, PT, SE 19970704 < 19970709 < | | | |
| R: DE, PRIORITY APPLN. | ES, FR, INFO.: | GB, IT | | JP 1996-180783 JP 1996-207011 | A 19960710 < A 19960806 < | | | |
| | | | | WO 1997-JP2378 | W 19970709 < | | | |

OTHER SOURCE(S): MARPAT 128:102004

GΙ

AB The title compds. [I; R1 = R4CO, R4CO, R4SO2, etc.; R2 = alkyl; R3 = H, acyl; R4 = (un)substituted C1-20 alkyl or C6-14 aryl, etc.; A = alkylene] are prepared I are efficacious in treating ischemic diseases, for example, ischemic brain diseases, cerebral stroke, cerebral thrombosis, cerebral embolism and myocardial infarction. Thus, compound (II; X = CO) (preparation

given) was reduced by (Me2CHCH2)2AlH to give 46% the title compound II (X = CHOH), which showed IC50 of 0.62 μM against calpain. ΤТ 167765-43-7P 201155-39-7P 201155-41-1P 201155-42-2P 201155-44-4P 201155-45-5P 201155-46-6P 201155-47-7P 201155-48-8P 201155-49-9P 201155-50-2P 201155-54-6P 201155-58-0P 201155-60-4P 201157-09-7P 201157-10-0P 201157-11-1P 201157-12-2P 201157-68-8P RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (preparation of hydroxytetrahydrofuran derivs. as remedies for ischemic diseases) 167765-43-7 CAPLUS RN CN Carbamic acid, [(1S)-1-[[[(2S,3S)-2-(acetyloxy)tetrahydro-3furanyl]amino]carbonyl]-3-methylbutyl]-, phenylmethyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 201155-39-7 CAPLUS
CN Carbamic acid, [(1S)-1-[[[(2S,3S)-2-(acetyloxy)tetrahydro-3-furanyl]amino]carbonyl]-3-methylbutyl]-, tetradecyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 201155-41-1 CAPLUS
CN Hexadecanamide, N-[(1S)-1-[[[(2S,3S)-2-(acetyloxy)tetrahydro-3-furanyl]amino]carbonyl]-3-methylbutyl]- (CA INDEX NAME)

Absolute stereochemistry.

RN 201155-42-2 CAPLUS
CN Benzamide, N-[(1S)-1-[[[(2S,3S)-2-(acetyloxy)tetrahydro-3-furanyl]amino]carbonyl]-3-methylbutyl]-2-fluoro- (CA INDEX NAME)

Absolute stereochemistry.

RN 201155-44-4 CAPLUS

CN Pentanamide, N-[(2S,3S)-2-(acetyloxy)tetrahydro-3-furanyl]-2-[(hexadecylsulfonyl)amino]-4-methyl-, (2S)- (CA INDEX NAME)

Absolute stereochemistry.

RN 201155-45-5 CAPLUS

CN Propanoic acid, 2,2-dimethyl-, (2S,3S)-tetrahydro-3-[[(2S)-4-methyl-1-oxo-2-[(phenylsulfonyl)amino]pentyl]amino]-2-furanyl ester (CA INDEX NAME)

RN 201155-46-6 CAPLUS

CN Benzamide, N-[(1S)-1-[[[(2S,3S)-2-(acetyloxy)tetrahydro-3-furanyl]amino]carbonyl]-3-methylbutyl]-4-chloro- (CA INDEX NAME)

Absolute stereochemistry.

RN 201155-47-7 CAPLUS

CN Pentanamide, N-[(2S,3S)-2-(acetyloxy)tetrahydro-3-furanyl]-4-methyl-2-[[(2,4,6-trimethylphenyl)sulfonyl]amino]-, (2S)- (CA INDEX NAME)

Absolute stereochemistry.

RN 201155-48-8 CAPLUS

CN Propanoic acid, 2,2-dimethyl-, (2S,3S)-3-[[(2S)-2-[[[4-(1,1-dimethylethyl)phenyl]sulfonyl]amino]-4-methyl-1-oxopentyl]amino]tetrahydro-2-furanyl ester (CA INDEX NAME)

RN 201155-49-9 CAPLUS

CN Pentanamide, N-[(2S,3S)-2-(acetyloxy)tetrahydro-3-furanyl]-2-[[(4-methoxyphenyl)sulfonyl]amino]-4-methyl-, (2S)- (CA INDEX NAME)

Absolute stereochemistry.

RN 201155-50-2 CAPLUS

CN Pentanamide, N-[(2S,3S)-2-(acetyloxy)tetrahydro-3-furanyl]-4-methyl-2-[(2-naphthalenylsulfonyl)amino]-, (2S)- (CA INDEX NAME)

Absolute stereochemistry.

RN 201155-54-6 CAPLUS

CN Carbamic acid, [(1S)-1-[[[(2S,3S)-2-(acetyloxy)tetrahydro-3-furanyl]amino]carbonyl]-3-methylbutyl]-, tridecyl ester (9CI) (CA INDEX NAME)

RN 201155-58-0 CAPLUS

CN Pentanamide, N-[(2S,3S)-2-(acetyloxy)tetrahydro-3-furanyl]-4-methyl-2-[(tetradecylsulfonyl)amino]-, (2S)- (CA INDEX NAME)

Absolute stereochemistry.

RN 201155-60-4 CAPLUS

CN Pentanamide, N-[(2S,3S)-2-(acetyloxy)tetrahydro-3-furanyl]-4-methyl-2-[(pentadecylsulfonyl)amino]-, (2S)- (CA INDEX NAME)

Absolute stereochemistry.

RN 201157-09-7 CAPLUS

CN Pentadecanamide, N-[(1S)-1-[[[(3S)-2-(acetyloxy)tetrahydro-3-furanyl]amino]carbonyl]-3-methylbutyl]- (CA INDEX NAME)

Absolute stereochemistry.

RN 201157-10-0 CAPLUS

CN Hexadecanamide, N-[(1S)-1-[[[(3S)-2-(acetyloxy)tetrahydro-3-furanyl]amino]carbonyl]-3-methylbutyl]- (CA INDEX NAME)

Absolute stereochemistry.

RN 201157-11-1 CAPLUS

CN Heptadecanamide, N-[(1S)-1-[[[(3S)-2-(acetyloxy)tetrahydro-3-furanyl]amino]carbonyl]-3-methylbutyl]- (CA INDEX NAME)

Absolute stereochemistry.

RN 201157-12-2 CAPLUS

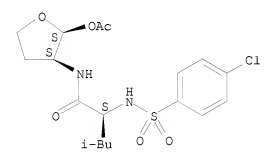
CN Pentanamide, N-[(2S,3S)-2-(acetyloxy)tetrahydro-3-furanyl]-4-methyl-2-[(phenylsulfonyl)amino]-, (2S)- (CA INDEX NAME)

Absolute stereochemistry.

RN 201157-68-8 CAPLUS

CN Pentanamide, N-[(2S,3S)-2-(acetyloxy)tetrahydro-3-furanyl]-2-[[(4-chlorophenyl)sulfonyl]amino]-4-methyl-, (2S)- (CA INDEX NAME)

Absolute stereochemistry.



REFERENCE COUNT: 6 THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 10 OF 10 CAPLUS COPYRIGHT 2009 ACS on STN

ACCESSION NUMBER: 1998:65807 CAPLUS

DOCUMENT NUMBER: 128:102386

ORIGINAL REFERENCE NO.: 128:20073a, 20076a

TITLE: Preparation and formulation of amino acid derivatives

for the prevention and treatment of neurodegenerative

diseases

INVENTOR(S): Yoshii, Narihiko; Saito, Ken-ichi; Ando, Ryoichi

PATENT ASSIGNEE(S): Mitsubishi Chemical Corp., Japan

CODEN: PIXXD2

SOURCE: PCT Int. Appl., 118 pp.

DOCUMENT TYPE: Patent

LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

| PA' | PATENT NO. | | | | KIND DATE | | APPLICATION NO. | | | Ο. | DATE | | | | |
|---------|------------|--------|-----|-----|-----------|-------|-----------------|-------|--------|---------|-------|-------|-------|-------|------|
| | | | | | | | | | | | | | | | |
| WO | 9801129 | 9 | | A1 | 1 | 19980 | 115 | WO | 1997- | -JP237 | 7 | | 1997 | 3709 | < |
| | W: US | 3 | | | | | | | | | | | | | |
| | RW: Al | Γ, BE, | CH, | DE, | DK, | ES, | FI, | FR, G | 3, GR, | IE, I | IT, I | LU, M | C, NL | , PT, | , SE |
| JP | 1010156 | 50 | | A | 1 | 19980 | 421 | JP | 1997- | -17975 | 7 | | 1997 | 3704 | < |
| PRIORIT | Y APPLN. | . INFO | .: | | | | | JP | 1996- | -180784 | 4 | A | 1996 | 3710 | < |

OTHER SOURCE(S):
GI

MARPAT 128:102386

AB The title compds. I [R1 represents R4CO, etc.; R4 represents alkyl, aryl or cycloalkyl; R2 represents alkyl; R3 represents hydrogen or acyl; and A represents alkylene] are prepared. These drugs are efficacious in preventing or treating neurodegenerative diseases, for example, Alzheimer's disease, diseases caused by demyelination in nerve cells, such as multiple sclerosis and neuropathy, and disorders accompanying cephalic traumas, such as consciousness disorder and motility disorder. $(3S)-3-((S)-4-Methyl-2-phenylsulfonylaminovalerylamino)-2-tetrahydrofuranol in vitro showed IC50 of 0.62 \ \mu M against calpain.$

Calpain.

167765-43-7P 201155-15-9P 201155-39-7P 201155-40-0P 201155-41-1P 201155-42-2P 201155-43-3P 201155-44-4P 201155-45-5P 201155-46-6P 201155-47-7P 201155-48-8P 201155-49-9P 201155-50-2P 201155-52-4P 201155-54-6P 201155-56-8P 201155-58-0P 201155-60-4P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of amino acid derivs. for prevention and treatment of neurodegenerative diseases)

RN 167765-43-7 CAPLUS

CN Carbamic acid, [(1S)-1-[[[(2S,3S)-2-(acetyloxy)tetrahydro-3-furanyl]amino]carbonyl]-3-methylbutyl]-, phenylmethyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 201155-15-9 CAPLUS

CN Pentanamide, N-[(3S)-2-(acetyloxy)tetrahydro-3-furanyl]-4-methyl-2-

[(phenylsulfonyl)amino]-, (2S)- (CA INDEX NAME)

Absolute stereochemistry.

RN 201155-39-7 CAPLUS

CN Carbamic acid, [(1S)-1-[[[(2S,3S)-2-(acetyloxy)tetrahydro-3-furanyl]amino]carbonyl]-3-methylbutyl]-, tetradecyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 201155-40-0 CAPLUS

CN Pentadecanamide, N-[(1S)-1-[[[(2S,3S)-2-(acetyloxy)tetrahydro-3-furanyl]amino]carbonyl]-3-methylbutyl]- (CA INDEX NAME)

RN 201155-41-1 CAPLUS

CN Hexadecanamide, N-[(1S)-1-[[[(2S,3S)-2-(acetyloxy)tetrahydro-3-furanyl]amino]carbonyl]-3-methylbutyl]- (CA INDEX NAME)

Absolute stereochemistry.

RN 201155-42-2 CAPLUS

CN Benzamide, N-[(1S)-1-[[[(2S,3S)-2-(acetyloxy)tetrahydro-3-furanyl]amino]carbonyl]-3-methylbutyl]-2-fluoro- (CA INDEX NAME)

Absolute stereochemistry.

RN 201155-43-3 CAPLUS

CN Benzamide, N-[(1S)-1-[[[(2S,3S)-2-(acetyloxy)tetrahydro-3-furanyl]amino]carbonyl]-3-methylbutyl]-4-fluoro- (CA INDEX NAME)

Absolute stereochemistry.

RN 201155-44-4 CAPLUS

CN Pentanamide, N-[(2S,3S)-2-(acetyloxy)tetrahydro-3-furanyl]-2-[(hexadecylsulfonyl)amino]-4-methyl-, (2S)- (CA INDEX NAME)

RN 201155-45-5 CAPLUS

CN Propanoic acid, 2,2-dimethyl-, (2S,3S)-tetrahydro-3-[[(2S)-4-methyl-1-oxo-2-[(phenylsulfonyl)amino]pentyl]amino]-2-furanyl ester (CA INDEX NAME)

Absolute stereochemistry.

RN 201155-46-6 CAPLUS

CN Benzamide, N-[(1S)-1-[[[(2S,3S)-2-(acetyloxy)tetrahydro-3-furanyl]amino]carbonyl]-3-methylbutyl]-4-chloro- (CA INDEX NAME)

Absolute stereochemistry.

RN 201155-47-7 CAPLUS

CN Pentanamide, N-[(2S,3S)-2-(acetyloxy)tetrahydro-3-furanyl]-4-methyl-2-[[(2,4,6-trimethylphenyl)sulfonyl]amino]-, (2S)- (CA INDEX NAME)

RN 201155-48-8 CAPLUS

CN Propanoic acid, 2,2-dimethyl-, (2S,3S)-3-[[(2S)-2-[[[4-(1,1-dimethylethyl)phenyl]sulfonyl]amino]-4-methyl-1-oxopentyl]amino]tetrahydro-2-furanyl ester (CA INDEX NAME)

Absolute stereochemistry.

RN 201155-49-9 CAPLUS

CN Pentanamide, N-[(2S,3S)-2-(acetyloxy)tetrahydro-3-furanyl]-2-[[(4-methoxyphenyl)sulfonyl]amino]-4-methyl-, (2S)- (CA INDEX NAME)

Absolute stereochemistry.

RN 201155-50-2 CAPLUS

CN Pentanamide, N-[(2S,3S)-2-(acetyloxy)tetrahydro-3-furanyl]-4-methyl-2-[(2-naphthalenylsulfonyl)amino]-, (2S)- (CA INDEX NAME)

RN 201155-52-4 CAPLUS

CN Heptadecanamide, N-[(1S)-1-[[[(2S,3S)-2-(acetyloxy)tetrahydro-3-furanyl]amino]carbonyl]-3-methylbutyl]- (CA INDEX NAME)

Absolute stereochemistry.

RN 201155-54-6 CAPLUS

CN Carbamic acid, [(1S)-1-[[[(2S,3S)-2-(acetyloxy)tetrahydro-3-furanyl]amino]carbonyl]-3-methylbutyl]-, tridecyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 201155-56-8 CAPLUS

CN Carbamic acid, [1-[[[2-(acetyloxy)tetrahydro-3-furanyl]amino]carbonyl]-3-methylbutyl]-, pentadecyl ester, [2S-[2α , 3α (R*)]]- (9CI) (CA INDEX NAME)

RN 201155-58-0 CAPLUS

CN Pentanamide, N-[(2S,3S)-2-(acetyloxy)tetrahydro-3-furanyl]-4-methyl-2-[(tetradecylsulfonyl)amino]-, (2S)- (CA INDEX NAME)

Absolute stereochemistry.

RN 201155-60-4 CAPLUS

CN Pentanamide, N-[(2S,3S)-2-(acetyloxy)tetrahydro-3-furanyl]-4-methyl-2-[(pentadecylsulfonyl)amino]-, (2S)- (CA INDEX NAME)

Absolute stereochemistry.

REFERENCE COUNT:

THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

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